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ACCESSION NUMBER: 92371258 EMBASE

DOCUMENT NUMBER:

1992371258

TITLE: Maitotoxin induces a calcium-dependent membrane

depolarization in GH4C1 pituitary cells via activation of

type L voltage-dependent calcium channels.

Xi D.; Van Dolah F.M.; Ramsdell J.S. AUTHOR:

CORPORATE SOURCE: Marine Biomedical/Environmental Sci., Medical University of

South Carolina, 221 Fort Johnson Rd., Charleston, SC 29412,

United States

Journal of Biological Chemistry, (1992) 267/35 SOURCE:

(25025-25031).

ISSN: 0021-9258 CODEN: JBCHA3

COUNTRY: United States

DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

water-soluble polyether, isolated from the marine dinoflagellate Gambierdiscus toxicus, that stimulates hormone release and Ca2+ influx. We have investigated the action by which MTX induces Ca2+ influx

and stimulates prolactin (PRL) release from GH4C1 rat pituitary cells. PRL release elicited by MTX is abolished in a concentration-dependent manner

by nimodipine, a dihydropyridine (DHP) antagonist of type L voltage- dependent calcium channels (L-VDCC), indicating that MTX-enhanced PRL release occurs via activation of type L-VDCC. As an

initial approach to. . . site. The effect of MTX on DHP binding was largely (65%) calcium-dependent. We next examined whether MTX alters the membrane potential of GH4C1 cells using the potential

sensitive fluorescent dye bisoxonol. Addition of 100

ng/ml MTX to GH4C1 cells caused a membrane depolarization within 2.5 min . . MTX-induced depolarization was not which reached a plateau. prevented by substitution of impermeant choline ions for Na+. It was similarly unaffected by K+ channel blockers or by depleting the K+ chemical concentration gradient with gramicidin, a monovalent cation pore-forming agent. By contrast, low extracellular Ca2+. component of the VDCC complex, which, in turn, initiates a positive feedback mechanism involving calcium-dependent membrane depolarization and

voltage-dependent activation of calcium channels.

ANSWER 15 OF 19 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.

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ACCESSION NUMBER: 92240513 EMBASE

DOCUMENT NUMBER: 1992240513

TITLE: Maitotoxin-induced intracellular calcium rise in PC12

cells: Involvement of dihydropyridine-sensitive and

ω-conotoxin-sensitive calcium channels and

phosphoinositide breakdown.

AUTHOR: Meucci O.; Grimaldi M.; Scorziello A.; Govoni S.;

Bergamaschi S.; Yasumoto T.; Schettini G.

CORPORATE SOURCE: Section of Pharmacology, Human Communicative Sciences

Dept., II School of Medicine, Via S. Pansini 5,80131

Napoli, Italy

SOURCE: Journal of Neurochemistry, (1992) 59/2 (679-688).

ISSN: 0022-3042 CODEN: JONRA

COUNTRY: United States DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 029 Clinical Biochemistry

> 052 Toxicology

LANGUAGE: English SUMMARY LANGUAGE: English

AB . calcium concentration and are always associated with an increase of the free cytosolic calcium level. We tested the effects of

voltage-sensitive calcium channel blockers

(nicardipine and ω -conotoxin) on maitotoxin-induced intracellular calcium increase, membrane depolarization, and inositol phosphate production in PC12 cells. Maitotoxin dose dependently. was reduced by pertussis toxin pretreatment. Maitotoxin caused a substantial membrane depolarization of PC12 cells as assessed by the fluorescent dye bisoxonol. This effect was reduced by pretreating the cells with either nicardipine or ω-conotoxin and was almost completely abolished by. in a calcium-free EGTA-containing medium. The findings on maitotoxin-induced cytosolic calcium rise and membrane depolarization suggest that maitotoxin exerts its action primarily through the activation of voltage-sensitive calcium channels, the increase of inositol phosphate production likely being an effect dependent on calcium influx. The ability of nicardipine and ω-conotoxin to inhibit the effect of maitotoxin on both calcium homeostasis and membrane potential suggests that L- and N-type calcium channel activation is responsible for the influx of calcium following exposure to maitotoxin,

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ACCESSION NUMBER: 92249069 EMBASE

DOCUMENT NUMBER:

1992249069

TITLE:

Membrane properties of identified mesencephalic dopamine

neurons in primary dissociated cell culture.

AUTHOR:

Chiodo L.A.; Kapatos G.

CORPORATE SOURCE:

1261 Scott Hall, Wayne State Univ. School of Medicine, 540

E. Canfield Ave., Detroit, MI 48201, United States

SOURCE:

Synapse, (1992) 11/4 (294-309). ISSN: 0887-4476 CODEN: SYNAET

United States

COUNTRY: DOCUMENT TYPE:

Journal; Article

FILE SEGMENT:

001 Anatomy, Anthropology, Embryology and Histology

002 Physiology

029 Clinical Biochemistry

LANGUAGE: English
SUMMARY LANGUAGE: English

SUMMARY LANGUAGE: English their distinct morphology, and this identification was validated with a double-labeling procedure that entailed the intracellular deposition of a fluorescent dye (Lucifer yellow or ethidium bromide), followed by processing for tyrosine hydroxylase immunocytochemistry. DA neurons identified in this manner were observed to have resting membrane potentials between -50 and -75 mV, input resistances of 50-360 M Ω , and membrane time constants of 4.1-14.1 msec. Forty-seven percent of. . . cells displayed spontaneous activity that was irregular in nature and often contained bursts (burst length was between two and six action potentials). The DA neurons displayed a variety of ionic conductances, including (1) a Na+ conductance (q(Na)) that underlies the action potential, (2) Ca2+ conductances (g(Ca)) that mediate the nonsomatic low- and high-threshold spikes observed, and (3) at least three K+ conductances (g(K)). Voltage-clamp analysis revealed several distinct transmembrane ionic currents, including (1) a large, rapidly inactivating tetrodotoxinsensitive inward Na+ current (I(Na)), (2) a 4-aminopyridinesensitive, transient early outward K+ current that required a conditioning hyperpolarization of the membrane to be activated by a subsequent depolarization. . . current was Ca2+-dependent and was not affected by tetraethylammonium ions. This current was termed I(AHP). The remaining current was not sensitive to changes in the extracellular Ca2+ concentration but was blocked by external tetraethylammonium. This current was termed I(K). The direct. (1-200 μM) onto the soma dose-dependently hyperpolarized these neurons; this effect was potentiated by the presence of the catecholamine reuptake blocker cocaine hydrochloride (10-200 µM). Under voltage-clamp conditions, DA was observed to increase I(K) significantly and had little effect on I(AHP). Thus, DA neurons in

monolayer cultures.

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ACCESSION NUMBER: 91049471 EMBASE

DOCUMENT NUMBER: 1991049471

TITLE: Bretylium causes a K+-Na+ pump activation that is

independent of Na+/H+ exchange in depolarized rat, mouse

and human lymphocytes.

AUTHOR: Tron L.; Pieri C.; Marian T.; Balkay L.; Emri M.;

Damjanovich S.

CORPORATE SOURCE: Biomedical Cyclotron Laboratory, University Medical School

of Debrecen, Debrecen, Hungary

SOURCE: Molecular Immunology, (1990) 27/12 (1307-1311).

ISSN: 0161-5890 CODEN: IMCHAZ

COUNTRY: United Kingdom

DOCUMENT TYPE: Journal; Conference Article

FILE SEGMENT: 026 Immunology, Serology and Transplantation

030 Pharmacology

037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

AB We have studied a bretylium tosylate induced increase of the membrane

potentials of partially depolarized rat, mouse and human

lymphocytes, using the **potential sensitive dye**, bis [1,3 dibutyl-barbituric acid-(5)- trimethine oxonol]. The extent of this depolarization is dose-dependent and decreased in magnitude as the temp was reduced from 37°C to room temperature The repolarizing effect is

inhibited by K+-Na+-pump blockers or lack of extracellular Na+.

Sodium ion channel **blockers** are effective in abolishing repolarization only if applied prior to, or simultaneously with,

bretylium. Activation of Na+/H+ exchange is not. . . is completely eliminated in the presence of 10 μ M amiloride (concn of the diuretics having no measurable inhibition on the **action** of the exchanger).

These data suggest that bretylium opens ligand- and voltage

-gated Na+ channels, and repolarization occurs due to higher activity of the K+-Na+-pump stimulated by the enhanced intracellular Na+ accumulation.

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COUNTRY:

ACCESSION NUMBER: 87120609 EMBASE

DOCUMENT NUMBER: 1987120609

TITLE: Maps of optical action potentials and NADH fluorescence in

intact working hearts.

AUTHOR: Salama G.; Lombardi R.; Elson J.

CORPORATE SOURCE: Department of Physiology, University of Pittburgh School of

Medicine, Pittsburgh, PA 15261, United States

SOURCE: American Journal of Physiology - Heart and Circulatory

Physiology, (1987) 252/2 (21/2) (H384-H394).

CODEN: AJPPDI United States

DOCUMENT TYPE: Journal

DOCUMENT TYPE: Journal

FILE SEGMENT: 002 Physiology

018 Cardiovascular Diseases and Cardiovascular Surgery

037 Drug Literature Index

LANGUAGE: English

AB Voltage-sensitive dyes were used to stain

intact perfused hearts and to simultaneously measure optical

action potentials (APs) from 124 sites on the

epicardium. Patterns of electrical depolarization (activation) and repolarization (recovery) along the surface of the. . . be altered by electrical stimulation. The normal heterogeneities in AP durations became

more pronounced in the presence of the Ca2+-entry blocker,

verapamil. The local metabolic state of the tissue was also monitored optically through its intrinsic NADH fluorescence measured from 124.

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ACCESSION NUMBER: 80026362 EMBASE

DOCUMENT NUMBER: 1980026362

TITLE: The effects of some organic 'calcium antagonists' on

calcium influx in presynaptic nerve terminals.

AUTHOR: Nachshen D.A.; Blaustein M.P.

CORPORATE SOURCE: Dept. Physiol. Biophys., Washington Univ. Med. Sch., St

Louis, Mo. 63110, United States

SOURCE: Molecular Pharmacology, (1979) 16/2 (579-586).

CODEN: MOPMA3 United States

COUNTRY: Un DOCUMENT TYPE: Jo

DOCUMENT TYPE: Journal

FILE SEGMENT: 037 Drug Literature Index

030 Pharmacology 002 Physiology

008 Neurology and Neurosurgery

LANGUAGE: English

The actions of the organic 'Ca antagonists' verapamil and D-600 were tested on pinched-off presynaptic nerve terminals (synaptosomes) from rat brain, and. . . or veratridine, an alkaloid that opens sodium channels. The extra uptake induced by depolarizing media appears to be mediated by voltage-sensitive Ca channels. Synaptosome depolarization was indirectly determined with the voltagesensitive fluorescent dye, di-pentyl oxacarbocyanine. Verapamil or D-600 (100 μM) inhibited the K+-induced 45Ca uptake by about two thirds, but had no effect. . . observations indicate that Na channels as well as Ca channels are inhibited by verapamil and D-600. Recordings of miniature end-plate potentials were used to evaluate the actions of verapamil and D-600 at the frog neuromuscular junction, after miniature end-plate potential frequency had been made sensitive to changes in the bathing Ca concentration by raising the external K+. Miniature end-plate potential frequency was not affected by verapamil (40-50 μM) or D-600 (10 µM) but was significantly reduced by Mn2+ (0.2 mM), a known blocker of Ca channels. Although verapamil and D-600 appear to be very potent antagonists of Ca currents in heart and smooth muscle, we conclude that Ca channels in vertebrate neurons are much less sensitive to these drugs.

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